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PRIME-BOOST VACCINATION STRATEGY

FIELD OF THE INVENTION:

The present invention relates to a method for inducing an immune

response to an antigen in a subject.

BACKGROUND OF THE INVENTION:

Measles is a highly contagious viral disease that has persisted for more Severe infection may lead to pneumonia, encephalitis (brain inflammation) than 1000 years since it was first described (Babbott and Gordon, 1954).

live-attenuated vaccine (LAV) it still causes approximately 800,000 deaths every year, predominantly among children in developing countries (Cutts and death. Although measles can be effectively prevented by a and Steinglass, 1998). 2

between 12 and 18 months. However maternal antibodies may decline more neutralization by maternal antibodies the LAV is generally administered consequence, there is a window between 8 and 18 months of age during neutralization of the vaccine by maternal antibodies. In order to avoid The inability to control measles using the LAV is largely due to rapidly in infants of developing countries (Gans et al., 1998). As which infants may lack both passive and active immunity.

requirement for trained staff for parenteral application of the vaccine, has led maintenance of the "cold chain" during transport and storage to ensure the An additional concern is the effective distribution and use of live ettenueted measles vaccines in developing countries in particular the viability of the vaccine prior to administration. This, together with

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titre Edmonston-Zagreb vaccine was given to young infants in the late 1980's. In an attempt to overcome the problem of maternal antibodies a high to poor vaccination coverage in these countries.

mortality from other infections such as diarrhoea and pneumonia (Markowitz in 1892 (Weiss, 1992). It is thought that the increase in mortality was due to et al., 1890; Garenne et al., 1991) and was subsequently withdrawn from use an immunosuppressive effect similar to that seen with wild type infection. This vaccine protected infants against measles but led to an increased ဓ္က

Sub-unit vaccines are not subject to the same constraints as LAVs. Development of a sub-unit vaccine for measles would primarily address

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(57) Abstract: The present invention provides a method for inducing an immune response to an amigen in a subject. The method i competies administrating to the subject DNA encoding the amigen, and subsequently orally administrating to the subject a composition comprising transgenic material, wherein the transgenic material comprises a DNA molecule encoding the antigen such that the antigen is expressed in the transgenic material.

(57) Abstract: The present invention provides a method for in Comparing administrating to the subject DNA encoding the subject DNA analogen is expressed in the transgenic material.

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Immuno-compromised pattents. New vaccine approaches such as DNA ssues concerning the immunization and protection of children in the developing world, such as maternal antibodies. In addition to this non-replicating sub-unit vaccines cannot initiate infection in

subunit vaccines and edible subunit vaccines are currently being devised as alternatives to the LAV. The measles virus (MV) hemagglutinin (H) protein is an immunodominant surface exposed glycoprotain and has been ncorporated into these vaccines.

encoding the MV-H protein. The immune responses generated have been of responses were not increased by multiple inoculations. In contrast, Yang et inoculation of BALB/c mice with a secreted form of plasmid DNA encoding BALB/c mice following repeated gene-gun inoculations. In addition, these the H protein induced a class I-restricted CTL response and IgG1 antibody were found to be 100-fold lower than those elicited by a single dose of the appropriate immune response on the number and route of administrations production (consistent with a $T_{\rm B}2$ -type response). Furthermore, antibody al. (1997) found that neutralizing antibody titres increased 2- to 4-fold in constructs were used for macaque vaccination, however, antibody levels varying success. Cardoso et al. (1996) demonstrated that intramuscular LAV (Polack et al., 2000). Such studies highlight the dependence of an tites were better than those raised by the LAV. When similar plasmid A number of studies have been conducted using DNA vaccines used in each particular animal model. 유 2

et al., 1898; Tacket et al., 1998; Wigdorovitz et al., 1999A & B). Systemic and their native immunogenic properties and are able to induce neutralizing and Bacterial and viral antigens have been expressed in transgenic plants protective antibodies in mice (Haq et al., 1995; Mason et al., 1998; Arakawa and transfently from plant viral vectors. Antigens from both sources retain mucosal immune responses have also been induced in human volunteers feed raw potato tubers expressing the binding subunit of the E. coll heat vaccination with plant-derived antigens can evoke a protective immune enterotoxin (LT) in vitro. Thus, the current data demonstrates that oral produced by these volunteers were able to neutralize E. coll heat labile abile enterotoxin (LT-B) (Tacket *et al.* 1998). The serum antibodies 23 8

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The present invention provides an alternate strategy for inducing an immune response to an antigen in a subject. Also provided are transgenic plants expressing an antigen derived from the measles virus.

SUMMARY OF THE INVENTION:

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In a first aspect, the present invention provides a method for inducing administaring to the subject DNA encoding the antigen, and subsequently sncoding the antigen such that the antigen is expressed in the transgenic orally administering to the subject a composition comprising transgenic an immune response to an antigen in a subject, the method comprising material, wherein the transgenic material comprises a DNA molecule material. 임

In a preferred embodiment of the present invention the composition further comprises a mucosal adjuvant, preferably cholera toxin β-subunits.

material as a fusion protein. In particular it is preferred the fusion protein It is also preferred that the antigen is expressed in the transgenic comprises the antigen C-terminally fused to the amino acid sequence SEXDEL (SEQ ID NO:1).

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The transgenic material is preferably a transgenic plant such as a fruit or vegetable. It is preferred that the transgenic plant is selected from the group consisting of; tobacco, lettuce, rice and bananas. 8

immunodeficiency virus, or Plasmodium sp. It is preferred that the antigen antigen is selected from the group consisting of viral antigens, parasitic is the measles virus H or F protein, or fragments thereof, preferably the In a further preferred embodiment of the present invention, the antigens and bacterial antigens, preferably measles virus, the human neasles H protein.

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In a still further preferred embodiment the DNA encoding the antigen is administered to the subject on at least two occasions and the composition comprising transgenic material is orally administered to the subject on at comprising transgenic material is orally administered to the subject on a least two occasions. More preferably, the DNA encoding the antigen is administered to the subject on a single occasion and the composition single occasion.

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the plant having been transformed with a DNA molecule, the DNA molecule In a second aspect the present invention provides a transgenic plant,

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comprising a sequence encoding a measles virus antigen such that the plant expresses the measles virus antigen.

In a preferred embodiment of this aspect of the invention, the DNA molecule encodes a fusion protein, preferably comprising the measles antigen C-terminally fused to the amino acid sequence SEKDEL.

In a further preferred embodiment the measles antigen is the measles protein.

Throughout this specification the word "comprise", or varietions such as "comprises" or "comprising", will be understood to imply the inclusion of a stated element, integer or step, or group of elements, integers or steps, but not the exclusion of any other element, integer or step, or group of elements, integers or steps.

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gene.

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The invention will hereinafter be described by way of the following non-limiting Figures and Examples.

BRIEF DESCRIPTION OF THE ACCOMPANYING DRAWINGS:

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Egure 1: Plant transformation vector constructs for expression of MV-H protein in tobacco. The T-DNA region inserted into the plant genome contains the nopaline synthese expression cassette [Kan*], which confers kanamycin resistance on transformed cells, and the MV-H protein expression cassette. The MV-H protein expression cassette comprises a cauliflower

(35S-Ter). The pBinH/KOEL and pBinSP/H/KDEL constructs contain an SEKDEL peptide sequence (KDEL) fused to the C-terminal end of the H protein for retention in the endoplasmic reticulum. The pBinSP/H/KDEL construct also contains a plant signal peptide (SP) fused to the N-terminal end of the H protein.

untranslated region (TEV) and cauliflower mosaic virus terminator sequence

mosaic virus 35S promoter (35S-Pro) fused to a tobacco etch virus 5'-

30 EXELTO 2: Transgene expression and production of recombinant MV-H protein in transgenic tobacco. (A) Northern blot comparing the level of MV-H gene expression of the six highest expressing T_o transgenic tobacco lines obtained for each MV-H construct. Each lane contained 10 µg of total RNA and was probed with a **P-labeled MV-H cDNA probe. (B) ELISA analysis of

MV-H protein expression in each of the T, transgenic tobacco lines shown in

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(A) detected with a rabbit anti-measles polyclonal antibody. Four

independent control transgenic lines transformed with a pBin construct lacking the MV-H gene, were included in analyses.

lacking the MV-H gene, were included in analyses.

Eigure 3: Detection of MV-H protein in pBinH/KDEL T, transgenic lines.

Selected kanamycin resistant progeny from the three highest T₀ expressing lines (8B, 12C and 39H) were analysed for MV-H protein expression using ELISA. The analysis was performed using either a rabbit anti-measles polyclonal antibody or MV-positive human serum. Control extract is from a transgenic tobacco line transformed with a pBin construct lacking the MV-H

Elgure 4. Immune response in mice following intraperitoneal (IP) immunization with transgenic plant extracts. Five mice were immunized with leaf extract from pBinHKDEL T, transgenic line 8B or a pBin control

- tansgenic line. IP immunizations were delivered on days 0, 14 and 49 with serum collected on days 28 and 84. (A) MV-specific serum IgG. Control serum is the mean value obtained from 3-4 naïve mice. (B) MV neutralization activity of serum IgG from day 84. MV-H (**), control (o).
- 20 Figure 5: Immune response in mice following gavage with transgenic plant extracts. (A) Mouse serum neutralization titres following gavage. Sera collected 49 days after initial treatment were pooled and the neutralizing ability against MV assessed in plaque-reduction neutralization (PRN) asseys. Neive (*), 28 MV-H + CT-CTB (*), and 28 control + CT-CTB (*).
 - 25 (B) MV-specific secretory IgA in fascal isolates collected 28 days after initial gavage.

Eigure 6: Serum MV neutralization (FRN) titres following DNA vaccination of mice. Sera collected 0, 15, 43 and 140 days after DNA vaccination were pooled. Natve (Φ), 2g MV-H + CT-CTB (Φ), and 2g control + CT-CTB (Ψ).

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Figure 7: MV-specific serum IgG titres following DNA-oral prime boost vaccination. Serum IgG titres were determined by ELISA on pooled sera from 0, 21 (pre-boost) and 49 days (post-boost). (A) MV-specific serum IgG titres for mice immunized with MV-H DNA and boosted with MV-H (-A-), or control (-F) plant extracts. (B) MV-specific serum IgG titres for mice

immunized with control DNA and boosted with MV-H (- Δ -), or control (- \blacksquare -) plant extracts. (C) Actual IgG titres represented in A and B.

Immunized with control DNA and boosted with MV-H (-▲-), or control (■-) MV-H (-▲-), or control (-■-) plant extracts. (B) Neutralization titre for mice Neutralization titre for mice immunized with MV-H DNA and boosted with Ekure 8: Serum MV neutralization (PRN) times following DNA-oral prime boost vaccination of mice. Neutralization titres were determined using plant extracts. (C) Actual neutralization titres represented in A and B. pooled sera from 0, 21 (pre-boost) and 49 days (post-boost). (A)

DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENTS:

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in the present invention are standard procedures, well known to those skilled Unless otherwise indicated, the recombinant DNA techniques utilized Cloning: A Practical Approach, Volumes 1-4, IRL Press (1995 and 1998); and F.M. Ausubel et al. (editors), Current Protocols in Molecular Biology, Greene Cloning, John Wiley and Sons (1984); J. Sambrook et al., Molecular Cloning: Brown (editor), Essential Molecular Biology: A Practical Approach, Volumes Pub. Associates and Wiley-Interscience (1988, including all updates until A Laboratory Manual, Cold Spring Harbour Laboratory Press (1989); T.A. in the art. Such techniques are described and explained throughout the 1 and 2, IRL Press (1991); D.M. Glover and B.D. Hames (editors), DNA literature in sources such as, J. Perbal, A Practical Guide to Molecular present) and are incorporated herein by reference.

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1996; Cox et al., 1993; Davis et al., 1993; Sedegah et al., 1994; Montgomery et encoding an antigen into tissues of a subject for expression of the antigen by al, 1993; Ulmer et al, 1993; Wang et al., 1993; Xiang et al., 1994; Yang et al., disclosures of which are hereby incorporated by reference in their entireties. vaccines" or "nucleic acid-based vaccines." DNA vaccines are described in experimental systems (see, for example, Conry et al., 1994; Cardoso et al., DNA vaccination involves the direct in vivo introduction of DNA the cells of the subject's tissue. Such vaccines are termed herein "DNA The ability of directly injected DNA that encodes an antigen to elicit a protective immune response has been demonstrated in numerous US 5,939,400, US 6,110,898, WO 95/20660 and WO 93/19183, the ဓ္က ន 8

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viral promoters derived from cytomegalovirus (CMV). These have had good To date, most DNA vaccines in mammalian systems have relied upon efficiency in both muscle and skin inoculation in a number of mammalian species. A factor known to affect the immune response elicited by DNA

plasmids, using a gene-gun, enhanced the immune responses of mice (Fynan immunization is the method of DNA delivery, for example, perenteral routes can yield low rates of gene transfer and produce considerable variability of efficiency of DNA transfection and more effective antigen presentation by transduction, cell fusion, DEAE dextran, calcium phosphate precipitation, gene expression (Montgomery et al., 1993). High-velocity inoculation of dendritic cells. Vectors containing the nucleic acid-based vaccine of the invention may also be introduced into the desired host by other methods et al., 1993; Eisenbraun et al., 1993), presumably because of a greater known in the art, e.g., transfection, electroporation, microinjection, 유

of biological origin that has been genetically engineered such that it produces "Transgenic material" of the present invention refers to any substance the antigen. Preferably, the transgenic material is a transgenic plant lipofection (lysosome fusion), or a DNA vector transporter.

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is used as a dietary component while the antigen is provided to the subject in consumption of a foodstuff, where the edible part of the transgenic material The orally administered composition can be administered by the ន

antigen in the DNA vaccine and/or the transgenic material but also allows for The present invention allows for the production of not only a single a plurality of antigens.

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thereby giving disease immunity even though the most prevalent strain is not organism. Alternatively, an organism may be sequentially or simultaneously transformed with a series of expression vectors, each of which contains DNA the expression of multiple antigenic amino acid sequences in one transgenic simultaneously boost an immune response to more than one of these strains, DNA sequences of multiple antigenic proteins can be included in the segments encoding one or more antigenic proteins. For example, there are expression vector used for transformation of an organism, thereby causing Transgenic material expressing multiple antigenic protein sequences can five or six different types of influenza, each requiring a different vaccine. cnown in advance. . ස

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part or in whole by a human or an animal such as, but not limited to, carrot, advantageous in certain disease prevention for human infants to produce a vaccine in a juice for ease of oral administration to humans such as tomato invention include any dicotyledon and monocotyledon which is edible in juice, soy bean milk, carrot juice, or a juice made from a variety of berry raspberries, banana and other such edible varieties. It is particularly Plants which are preferably used in the practice of the present potato, apple, soybean, rice, corn, berries such as strawberries and

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types. Other foodstuffs for easy consumption include dried fruit.

- as well. Such tissue would include but would not be limited to embryogenic Several techniques extet for introducing foreign genetic material into a coated onto microparticles directly into cells (see, for example, US 4,945,050 technology (see, for example, US 5,177,010, US 5,104,310, US 5,004,863, US plants, the type of tissue which is contacted with the foreign genes may vary tissue, callus tissue type I and II, hypocotyl, meristem, and the like. Almost herein by reference. In addition to numerous technologies for transforming Introduced gene. Such techniques include acceleration of genetic material plant cell, and for obtaining plants that stably maintain and express the 5,159,135). Electroporation technology has also been used to transform 92/09898 and WO 93/21335). Each of these references are incorporated and US 5,141,131). Plants may be transformed using Agrobacterium plants (see, for example, WO 87/08614, US 5,472,869, 5,384,253, WO all plant tissues may be transformed during development and/or differentiation using appropriate techniques described herein. 2 15 2
 - A number of vectors suitable for stable transfection of plant cells or for and Weissbach, Methods for Plant Molecular Biology, Academic Press, 1989; one or more cloned plant genes under the transcriptional control of 5' and 3' the establishment of transgenic plants have been described in, e.g., Pouwels et al., Cloning Vectors: A Laboratory Manual, 1985, supp. 1987; Weissbach regulatory region controlling inducible or constitutive, environmentally- or Publishers, 1990. Typically, plant expression vectors include, for example, expression vectors also can contain a promoter regulatory region (e.g., a and Gelvin et al., Plant Molecular Biology Manual, Kluwer Academic regulatory sequences and a dominant selectable marker. Such plant developmentally-regulated, or cell- or tissue-specific expression), a ဓ္ဗ 33 23

ranscription initiation start site, a ribosome binding site, an

RNA processing signal, a transcription termination site, and/or a polyadenylation signal.

Examples of plant promoters include, but are not limited to ribulosepromoter, phaseolin promoter, ADH promoter, heat-shock promoters and tissue specific promoters. Promoters may also contain certain enhancer 1,8-bisphosphate carboxylase small subunit, beta-conglycinin

sequence elements that may improve the transcription efficiency. Typical

enhancers include but are not limited to Adh-intron 1 and Adh-intron 6.

promoters are responsible for gene expression in specific cell or tissue types, such as the leaves or seeds (e.g., zein, oleosin, napin, ACP, globulin and the tissues and organs. Examples of such promoters include but are not limited Constitutive promoters direct continuous gene expression in all cells like) and these promoters may also be used. Promoters may also be active to pollen-specific, embryo specific, corn silk specific, cotton fiber specific, during a certain stage of the plants' development as well as active in plant types and at all times (e.g., actin, ubiquitin, CaMV 35S). Tissue specific root specific, seed endosperm specific promoters and the like. 9 12

desirable transcription and translation elements that function in plants may response to a specific signal, such as: physical stimulus (heat shock genes); promoter. An inducible promoter is responsible for expression of genes in Under certain circumstances it may be desirable to use an inducible light (RUBP carboxylase); hormone (Em); metabolites; and stress. Other be used.

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nopaline synthase promoter, the mannopine synthase promoter; promoters of In addition to plant promoters, promoters from a variety of sources can viral origin, such as the cauliflower mosaic virus (35S and 19S) and the like promoters of bacterial origin, such as the octopine synthase promoter, the be used efficiently in plant cells to express foreign genes. For example, may be used. 23

1998; Kapustra et al., 1999; Brennan et al., 1999). It has been suggested that the particulate form of these VLPs or chimeric viruses may result in greater transgenic plants producing virus-like particles (VLPs), or chimeric plant viruses displaying antigenic epitopes (Mason et al., 1996; Modelska et al., developed for both animal and human pathogens (Hood and Jilka, 1999). A number of plant-derived edible vaccines are currently being Immune responses have also resulted from oral immunization with 33 ဓ္က

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tability of the antigen in the stomach, effectively increasing the amount of antigen avallable for uptake in the gut (Mason et al. 1998, Modelska et al.

as to change one or more amino acid residues in the antigen expressed in the portion of an antigenic protein as either a smaller peptide or as a component expression vectors may contain DNA coding sequences which are altered so protein. Expression vectors containing a DNA sequence encoding only a particular antigen may also be utilized in this invention. For example, Mutant and variant forms of the DNA sequences encoding for a transgenic material, thereby altering the antigenicity of the expressed of a new chimeric fusion protein are also included in this invention.

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proper immunization of pets. Viral vaccines for diseases such as: Newcastle, The present invention can be used to produce an immune response in production animals such as: brucellosis, fowl cholera, anthrax and black leg material used in the methods of the present invention may be incorporated canine hepatits, parvorirus, and feline leukemia may be controlled with Rinderpest, hog cholera, blue tongue and foot-mouth can control disease economic losses from disease deaths. Prevention of bacterial diseases in animals other than humans. Diseases such as: canine distemper, rabies, through the use of vaccines has existed for many years. The transgenic outbreaks in production animal populations, thereby avoiding large into the feed of animals.

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A "mucosal adjuvant" is a compound which non-specifically stimulates or enhances a mucosal immune response (e.g., production of IgA antibodies). induction of a mucosal immune response to the immunogenic compound. Administration of a mucosal adjuvant in a composition facilitates the

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The mucosel adjuvent may be any mucosel adjuvent known in the art adjuvant may be cholera toxin (CT), enterotoxigenic B. Coll heat-labile toxin idjuvanticity. Preferably, the mucosal adjuvant is cholera toxin B-subunits. The mucosal adjuvant is co-administered with the composition comprising immune response. The suitable amount of adjuvant may be determined by which is appropriate for human or animal use. For example, the mucosal transgenic material in an amount effective to elicit or enhance a mucosal standard methods by one skilled in the art. Preferably, the adjuvant is (LT), or a derivative, subunit, or fragment of CT or LT which retains

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present at a ratio of 1 part adjuvant to 10 parts composition comprising the

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ransgenic material

In the present invention, the antigen can be expressed in the

- same or different native protein. In the latter instance, the different antigens ransgenic material as a fusion protein. Typically, the additional amino acid also preferred that the fusion protein comprise at least two antigens from the antigen. Preferably, the fusion protein results in a higher immune response when compared to when the antigen not expressed as a fusion protein. It is can be from different organisms, providing immune protection against a sequence will extend from the C-terminus and/or the N-terminus of the ß 유

number of pathogens.

Experimental Protocol

Construction of transgenic tobacco plants producing H protein 12

Three constructs were generated for the expression of MV-H protein in (c) pBinSP/H/KDEL - addition of both an N-terminal plant signal peptide and addition of a C-terminal endoplasmic reticulum (ER)-retention sequence and tobacco plants (Figure 1) (a) pBinH – H protein alone, (b) pBinH/KDEL – a C-terminal ER-retantion sequence.

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Hopkins Hospital, Baltimore). Using the Altered Sites kit (Promega) an Nool GenBank accession no. X16565) was obtained from plasmid pBS-HA (Johns encompassing the open reading frame of the MV-H gene (Edmonston strain; To produce these constructs a 1.8 kb EcoRI / BanHI fragment

- protein from serine to alanine. The Nool / BamHI fragment containing the Nsecond codon from T to C. This also altered the second amino acid of the H site was introduced into the 5'-end of the H gene. The Ncol site was created around the existing initiation codon by mutating the first nucleotide of the terminal modified H gene was then transferred into the plant expression 23
 - A second H-protein construct containing the Ncol site described above terminus of the H gene immediately upstream of the stop codon and BamHI Pelham, 1987) was also engineered. A XhoI site was introduced into the Cand an andoplasmic raticulum-ratantion sequence SEXDEL (Munro and vector pRTL2 (Restrepo et al., 1990) to give pRTL2-H. ဓ္က
 - site using the Altered Sites Ltt (Promega). This allowed a double-stranded oligonucleotide encoding the SEKDEL sequence to be ligated between the 33

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Xhol and BamHI sites creating an in-frame fusion with the C-terminal end of the H protein. The SEKDEL oligonucleotide was produced by annealing the following complementary sequences: 5'-

- D NO:4) to TNLQSEXDEL* (SEQ ID NO: 5). The HXDEL fragment was then terminal sequence of the modified H protain was altered from TNRR* (SEQ GATCCCCTCATAGCTCAT CTTTCTCAGAGA-3' (SEQ ID NO.3). The C-TCGATCTCTGAGAAGATGAGCTATGAGGG-3' (SEQ ID NO:2) and 5'cloned into pRTL2 to give pRTL2-H/KDEL.
- In the third construct, the signal peptide (SP) of the tobacco Pria gene (Hammond-Kosack et al. 1994) was cloned into the Nool site of pRTL2-HKOEL upstream of, and in frame with, the H protein. The 107 bp SP fragment was amplified by PCR from the plasmid SLJ6069 (Sainsbury Laboratory, JIC, Norwich, UK) using the oligonucleotides: 5'-GCCCCATGCCATTTGTTCTTTT-3' (SEQ ID NO: 8) and 5'-유
- TATCCATGGGCCCGGCACGCCAAGAGTGGGATAT-3! (SEQ ID NO:7). This clone was designated pRTL2-SP/H/KDEL. 12

expression cassettes of pRTL2-H, pRTL2-H/KDEL, and pRTL2-SP/H/KDEL were transferred into the binary vector pBin19 (Bevan, 1984) to produce Pollowing verification of modifications by sequence analysis, the pBinH, pBinH/KOEL and pBinSP/H/KOEL, respectively (Figure 1).

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(Nicotlana tabacum var Samsun) using the leaf disc method as described by These three constructs were then electroporated into Agrobacterium tumefaciens strain LBA 4404 and used for transformation of tobacco Horsch et al. (1985).

Transgene expression analysis

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The pellet was rinsed with cold 70% ethanol, dried and resuspended in 25 µl (25:24:1 v/v). The final aqueous phase was mixed with 0.1 volume of sodium for 30 min and nucleic acid pelleted by centrifugation at 13,000 g for 10 min. of sterile water. RNA was analysed by northern blot using a *P-labelled MV-1% 9-mercaptoethanol, pH 9.0 by extracting twice with an equal volume of acetate (pH 5.0) and 2.5 volumes of cold 100% ethanol, incubated at -20°C transganic tobacco plants in 0.1M Tris, 0.1M NaCl, 10 mM EDTA, 1% SDS, phenol and once with equal volume of phenol:chloroform:isoamyl alcohol Total RNA was extracted from 150mg leaf samples of in vitro

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Detection of MV-H protein in transgenic tobacco by EUSA

Tobacco leaves (50mg) were frozen in liquid nitrogen and ground to a fine powder in a 1.5 ml eppendorf. Five volumes of chilled extraction buffer extract was then centrifuged at 23,000 g for 15 min at 4°C, the supernatant (PBS containing 100mM ascorbic acid, 20mM EDTA, 0.196 Tween-20 and collected and glycerol added to a final concentration of 18% before snap 1mM PMSF, pH 7.4) was added and the extract vortexed for 15 s. The freezing in liquid nitrogen and storage at -70°C.

- Plant extracts were diluted in 0.1M carbonate buffer (pH 9.8) and were coated onto ELISA plates at 4°C overnight. All further incubations were at 37°C for 1 hour. Pollowing a blocking step with 2.5% skim milk the MV-H protein was detected with a rabbit polyclonal anti-measles antibody (CDC, Atlanta) diluted 1/4000. Anti-rabbit horseradish peroxidase conjugate 2
 - (Boshringer Mannheim) diluted 1/8000 was used as the secondary antibody. The plates were developed with TMB (3,3,5,5'-tetramethylbenzidine) substrate for 30 - 60 min and read at 630nm.

Preparation of antigen from transgenic plants

- performed on ice or at 4°C. Frozen tobacco leaves were powdered in a coffee Glycerol was added to the pellet to a final concentration of 16% allowing the above). The extract was filtered through 2 layers of miracloth, centrifuged at extracts to be stored at -70°C. Extracts ranged in concentration from 3.2g/ml 100g for 5 min and the supernatant centrifuged again at 32,600 g for 60 min. grinder and mixed with 2.5 volumes of chilled extraction buffer (described pBinH/KDEL transgenic line 8B, or transgenic tobacco lacking the MV-H Recently expanded leaves from glasshouse grown plants of the gene, were harvested and stored at -35°C. All subsequent steps were to 4.5g/ml. 22 2
- The supernatant from the 32,800g spin was further purified. Proteins containing 10 mM ascorbic acid, and applied to PD-10 columns (Amersham Pharmacia Biotech, Uppsala, Sweden) pre-equilibrated with PBS. The precipitated from the supernatant between 25% and 50% ammonium sulphate (AS) were resuspended in phosphate buffered saline (PBS) protein fraction was eluted in PBS, glycerol was added to a final 35 8

concentration of 18% allowing the extracts to be stored at -70°C.

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A mucosal adjuvant consisting of 2µg of cholera toxin (CT) and 10µg of needle attached to a 1ml Tuberculin syringe. The gavage needle was inserted down the oesophagus of anaesthetized animals into the stomach, where 0:4g, immediately prior to gavage. Gavage was performed using an 8cm gavage 18, 28 or 48 of plant material was injected. Mice were studied for signs of cholera toxin B subunit (CTB) (Sigma, USA) was added to plant aliquots tracheal or nasal obstruction until fully recovered from anaesthetic.

Laboratory mice and cell lines

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old), were purchased from Animal Research Centre, Western Australia, and cells (RMK cells) were grown as monolayers at 37°C in RPMI 1640 medium (Trace, Biosciences Ltd, Australia) supplemented with 10% fetal calf serum were maintained in the University Animal House, Rhesus monkey kidney Adult female Balb/c mice, between 18-25g (approximately 8 weeks (FCS) (Trace) in a 5% CO, atmosphere

Construction and vaccination of MV-H DNA

12

containing the extracellular domain of the measles virus H gene (MV-H), and ampicillin resistance and the SV40 late polyadenylation signal was used for A high copy pCI plasmid vector (Promega, USA) incorporating a vaccine production. Two DNA vaccine constructs were prepared. One human cytomegalovirus (CMV) immediate-late enhances/promoter, a control construct containing the ovalburnin gene.

A 1ml Insulin needle (Becton Dickinson, USA) was used to inject 25 or 50µg of DNA solution into both quadriceps of each mouse.

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Collection of mouse samples

Blood was collected by intraocular bleeding or cardiac puncture, once blood had clotted serum was recovered by centrifugation (7100g, 6 min).

- Paeces were collected into eppendorfs pre-blocked with 1% BSA. 1ml by vortexing then centrifuged (25,000g, 6 min). The supernatant was stored faecas. Following overnight incubation at 4°C, solid material was disrupted of 0.1% BSA + 0.15mM PMSF solution in PBS was added per 100mg of at -20°C in pre-blocked eppendorfs. 8
- To collect saliva samples anaesthetized mice were injected with 200µl of 20µg/ml carbachol in PBS to induce salivation. 33

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serum in PBS) was inserted. After dispensing wash solution into the lungs, a region was exposed and muscle tissue surrounding the trachea removed. A plunger and the extraction of lung fluid. Two more washes were performed 10 second rib-cage massage was performed prior to retraction of the syringe Tuberculin syringe containing 0.4ml of wash solution (1% v/v foetal calf Bronchoalveolar fluid was collected from killed mice. The throat small hole was made in the traches and a lavage tip attached to a 1ml using 0.3ml of wash solution.

Detection of MV-specific antibodies 10

Enzygnost measles-coated plates (Dade-Behring, Germany), containing peroxidase-conjugated goat anti-mouse IgG followed by tetramethyl-bromide simian kidney cells infected with MV, were used for detection of anti-MV antbody in mouse samples. MV-specific antbodies were detected with (TMB) substrate. IgG-typing was performed using alkaline phosphatase (AP) -conjugated anti-mouse IgG1 or AP-conjugated anti-mouse IgG2a and p-Nitrophenyl phosphate (pNPP) substrate.

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Mouse serum, salivary, BAL and faecal samples were assayed for the presence of IgA using AP-conjugated goat anti-mouse IgA with pNPP

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Plague reduction neutralization assay

The plaque reduction neutralization (PRN) titre is the reciprocal of the serum dilution capable of preventing 50% plaque formation by wild-type MV. The Edmonston strain of MV was used for this assay.

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(200pfu/100µl). This serum/virus suspension was incubated at 37°C for 90 supplemented RPMI (1/4 to 1/4098) and added to an equal volume of MV Four-fold dilutions of heat inactivated sera were prepared in

confluent RMK cells. Following a 90 minute incubation at 37°C 1ml/well of supplemented RPMI medium was added and plates were incubated at 37°C minutes before addition to 24-well, flat-bottomed plates containing 80% in a humidified atmosphere of 5% CO₂ for 72 hours. 9

Growth medium was removed and cells were fixed and permeabilised with 1ml/well of 10% formaldehyde with 0.1% Triton-X 100 in PBS for 20 minutes at RT. Plates were blocked with goat serum and anti-MV IgG 33

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a Leitz fluovert inverted fluorescent microscope. Each cluster of fluorescing, FITC-conjugated anti-human IgG and fluorescing cells were examined using positive human serum was added. Anti-MV human sera was detected with preventing 50% plaque formation was generated according to the Karber infected cells was counted as one pfu. The serum dilution capable of formula.

Results

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Transgenic tobacco plants producing MV-H protein

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targeting on antigen yield, two additional clones were constructed, with a C-Munro and Pelham 1987), and an authentic N-terminal plant signal peptide terminal SEKDEL sequence, coding for retention in the ER (pBinH/KDEL; expression cassette (Figure 1). To compare the effect of intracellular hemagglutinin (H) gene (Edmonston strain) was cloned into a plant A 1.8kb fragment encompassing the coding region of the MV

A total of 90 primary transformant (T₀) lines were obtained which [pBinSP/H/KDEL; Hammond-Kosack et al., 1994].

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rabbit anti-messles polyclonal antibody (Figure 2B). Plants transformed with showed detectable levels of MV-H gene expression by northern blot analysis (data not shown). A comparison of the six highest expressing lines for each construct are shown in Figure 2A. Transgene expression was similar for all further analysed for level of recombinant MV-H protein by ELISA using a three constructs. The selected high expressors shown in Figure 2A were the pBinH construct produced small quantities of recombinant MV-H 20

lines containing constructs pBinH and pBinH/KDEL, there appeared to be a pBinSP/HKDEL lines relative to the HKDEL transgenic lines. For tobacco protein. However, addition of the C-terminal KDEL sequence resulted in with the pBinH/KDEL construct. Interestingly, addition of the Pria plant much higher levels of MV-H protein accumulation in plants transformed signal peptide appeared to inhibit MV-H protein production in 23 8

Seed was collected from the pBinH/KDEL To transgenic lines showing kanamycin and re-assayed for MV-H protein production. ELISA analysis using the rabbit and-measles polyclonal andserum showed that the the highest levels of H production (12C, 8B & 39H), germinated on

reasonable correlation between transgene expression level and MV-H protein

production (compare Figures 2A & 2B).

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introduced MV-H transgene was stably inherited in the T₁ progeny (Figure 3). measles polyclonal antiserum (Figure 3), confirming that the plant-derived MV-H protein retained at least some of the antigenic regions present in the obtained from a subject with a history of wild-type measles infection, who detected similar quantities of MV-H protein in T, plants as the rabbit antihad tested positive for measles antibodies by ELISA. The human serum pBinH/KDEL T, progeny by human serum (Figure 3). This serum was Recombinant MV-H protein could also be detected in leaf extracts of native MV-H protein.

monoclonal antibodies as tested by indirect ELISA. MAb-386 detected MV-H values ranging from 0.063 to 0.065 for the pBinH/XDEL extracts, compared to Further evidence of the authentic antigenicity of the recombinant MVranging from 0.392 to 0.420, compared to 0.018 to 0.019 for extracts from pBin control transgenic. The response of MAb-CV4 provided absorbance protein in extracts of pBinH/KOEL 8B (T.) line with absorbance readings H protein was its positive reaction with two out of three MV-H protein -0.005 to -0.001 for control transgenic extracts. 13 9

Intraperitoneal vaccination with plant-derived MV-H protein induces MV neutralizing antibodies

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on day 0, 14 and 49 and serum was collected on day 28 and 84. Significantly plant extract from MV-H or control transgenic plants. Mice were inoculated To determine the immunogenicity of the plant-derived MV-H protein (Figure 4B). These results demonstrate that plant-derived MV-H protein is groups of BALB/c mics were inoculated intraperitoneally with AS-purified MV-H than in mice inoculated with control plant extract (P < 0.01) (Figure more MV-specific IgG was detected in mice vaccinated with plant-derived 4A). The MV-specific IgG was able to neutralize wild-type MV in vitro ímmunogenic when administered intraperitoneally. 엃

Oral vaccination with plant-derived MV-H protein induces, neutralizing antibodies and sigA

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have developed neutralizing antbodies to wild-type MV, detalls of one of Mice gavaged with either AS-purified MV-H or pellet MV-H extract these expertments are given below.

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containing the mucosal adjuvant CT-CTB by gavage on days 0, 7, 14, 21 and 35. Sera were collected on days 0, 7, 14, 21, 28, 49 and 78 and faecal isolates but for only 49 days in mice gavaged with 4g of extract, with maximum titres responses persisted for at least 78 days in mice gavaged with 2g of extract, obtained on days 0 and 28. MV-specific serum IgG was only detected in of 2187 and 9 respectively. The lower response to 4g may be due to the groups that received 2g or 4g of MV-H plant extract. The serum IgG Groups of three mice were given 1g, 2g or 4g of plant extract increased dose to tobacco toxins also received.

mice gavaged with 2g of MV-H plant extract (Figure 5A). It peaked at 78 days with a PRN titre of 600. Mice gavaged with 4g of MV-H plant extract had a maxdmum neutralization titre of 150 at day 49. No neutralizing ability was High neutralizing ability was observed in pooled sera collected from detected in mice gavaged with 2g of control plant extract. 2

particularly important result as mucosal immunity is the first line of defense MV-specific secretory IgA (sIgA) was detected in faecal samples from some mice gavaged with 2g of MV-H plant extract (Figure 5B). This is a against airborne pathogens such as measles.

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ovalbumin DNA (control) on day 0. Sera was collected on days 0, 15, 43 and experimental group that received MV-H DNA. High serum IgG levels were Vaccination with MV-H DNA constructs induces MV-neutralizing antibodies after vaccination an increase in MV-specific IgG was only observed in the 140, and faecal samples were obtained on days 0, 7, 14 and 21. Ten days to mice immunized with control DNA, which produced no MV-specific Groups of five mice were injected with 100µg of MV-H DNA, or . 8 23

immune response, serum IgG from mice primed with MV-H DNA was able to recorded at day 140, suggesting that the immune response is persistent. High thes of MV-neutralizing antibodies have previously been raised using MV-H neutralize wild-type MV in vitro (Figure 6). A neutralization titre of 900 was maintained from day 20 to day 43, with a maximum titre of 729. In contrast DNA vaccines in mice (Yang et al. 1997, Polack et al. 2000), however some studies suggest that maternal antibodies many interfere with vaccine efficiency (Schlereth et al. 2000). ဓ္က

The predominant isotype present in mice immunized with MV-H DNA was IgG1, indicating a T_R2-type response. While intramuscular DNA 33

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hemagglutinin-based influenza DNA vaccine (Cardoso *et al.* 1996, Deliyannis vaccines are generally associated with $T_{\rm H}1$ -type responses, $T_{\rm H}2$ dominated from, rather than retained within, transfected cells, although there are no responses have been reported to occur in response to intramuscular DNA difference in antigen presentation when the encoded antigen is released et al. 2000). It is possible that this switching of IgG isotypes is due to a vaccination with a secreted form of measles H protein and a secreted conclusive data to account for these differences.

No MV-specific serum or secretory IgA was detected in any DNA immunized group.

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Oral delivery of MV-H protein following MV-H DNA vaccine boosts serum IgG

days 0, 21 (pre-boost), and 49 (post-boost), and faecal isolates were obtained days 21, 28, 35 and 42, these mice were boosted with 2g of either control or H protein plant extract, administered with CT-CTB. Sera were collected on Mice were primed with 50µg of MV-H or control DNA on day 0. On weekly until day 49. Salivary and bronchoalveolar lavage (BAL) samples were collected on day 49. Five mice were used per treatment. 5

MV-specific IgG, but mice given control DNA did not. The titre of the MV-H serum IgG in mics primed with control DNA and boosted with control plant extract indicates that this is due to a continuing response to the MV-H DNA post-boost pooled sera (Figure 7). Mice primed with MV-H DNA, produced raccine followed by an oral MV-H plant boost resulted in higher serum IgG plant extract. MV-H DNA primed mice boosted with control plant extracts ulso had higher post-boost IgG titres. However the absence of MV-specific DNA IgG response was increased three-fold following gavage with MV.H raccine and not to the control plant extract. Delivery of the MV-H DNA MV-specific serum IgG times were determined for pre-boost and ន ĸ

Oral delivery of MV-H protein following MV-H DNA vaccine boosts neutralization titres

thes than either DNA vaccination or oral plant vaccination alone (MV-H

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INA-control plant and control DNA - MV-H plant respectively),

Neutralization assays were performed on pooled sera collected prior to DNA vaccination (day 0), immediately before boosting with plant extracts 33

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(day 21) and 1 week after the final plant feeding (day 49) for each of the four treatment groups.

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(Figure 8). At day 21 (pre-boost) serum from MV-H DNA primed mice had an plant extract (Figure 8). The neutralization titre for MV-H DNA primed mice neutralization titres increased relative to titres for mice boosted with control average neutralization titre of 1150 compared to a titre of 8 for mice primed boosted with control plant dropped from 1150 to 450, while mice boosted The neutralization titres exhibited similar trends to the IgG titres with control DNA. Following gavage with MV-H plant extracts

As with serum IgG titres combining the MV-H DNA vaccine and MV-H plant extract resulted in a synergistic response producing neutralization tites in excess of those recorded for either DNA or plant extract alone

anhanced both the magnitude and the persistence of the immune response.

with MV-H plant extract exhibited an increase in neutralization titre from

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1150 to 2550. This suggests that boosting with MV-H plant extract has

(Figure 8). 12

Finally, oral immunization using plant-derived MV-H protein resulted in the infection. Furthermore the present invention shows that mice immunized Neutralization titres for serum IgG were greater following DNA-oral prime recognised by host antibodies produced in response to wild-type measles boost than when either DNA or plant extracts were used alone (Figure 8). intraperitoneally, by gavage or by DNA-oral prime-boost all developed antbodies able to neutralize wild-type MV in vitro (Figures 4B, 5A, 8). expressed in transgenic material and that this recombinant protein is The present invention demonstrates that MV-H protein can be production of measurable levels of MV-specific sigA (Figure 5B).

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important advances of this vaccine strategy. Availability of the vaccine in an vaccination coverage by providing an inexpensive and relatively heat-stable "edible" form as a constituent of a fruit or vegetable crop will also enhance prima-boost" vaccination strategy utilising transgenic organisms is a viable approach to new vaccines. The potential for inducing a mucosal immune package for distribution. Such a vaccine will have the potential to enable response, and seroconversion in the presence of maternal antibodies are rates of vaccination to reach the targets required for global eradication. The present study demonstrates that "DNA vaccination-oral ജ æ

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invantion as broadly described. The present embodiments are, therefore, to variations and/or modifications may be made to the invention as shown in the specific embodiments without departing from the spirit or scope of the It will be appreciated by persons skilled in the art that numerous be considered in all respects as illustrative and not restrictive.

purpose of providing a context for the present invention. It is not to be taken as an admission that any or all of these matters form part of the prior art base Any discussion of documents, acts, materials, devices, articles or the invention as it existed in Australia before the priority date of each claim of ilke which has been included in the present specification is solely for the or were common general knowledge in the field relevant to the present this application. 유

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- A method for inducing an immune response to an antigen in a subject, andgen, and subsequently orally administering to the subject a composition the method comprising administering to the subject DNA encoding the
 - DNA molecule encoding the antigen such that the antigen is expressed in the comprising transgenic material, wherein the transgenic material comprises a ransgenic material.
 - A method as claimed in claim 1 in which the composition further comprises a mucosal adjuvant. ٠i 유
- A method as claimed in claim 2 in which the mucosal adjuvant is က်
 - cholera toxta β-subunits.

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- A method as claimed in any one of claims 1 to 3 in which the antigen is expressed in the transgenic material as a fusion protein.
- A method as claimed in claim 4 in which the fusion protein comprises the antigen C-terminally fused to the amino acid sequence SEKDEL ន
- A method as claimed in any one of claims 1 to 5 in which the transgenic material is a transgenic plant. ø.
- A method as claimed in claim 8 in which the transgenic plant is a fruit or vegetable. ۲. 23
- selected from the group consisting of; tobacco, lettuce, rice and bananas. A method as claimed in claim 6 in which the transgenic plant is œ
- is selected from the group consisting of viral antigens, parasitic antigens and A method as claimed in any one of claims 1 to 8 in which the antigen bacterial antigens.

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10. A method as claimed in claim 9 in the which the antigen is from measles virus, the human immunodeficiency virus, or Plasmodium sp. 33

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- the group consisting of the measles virus H or F protein, or fragments thereof. 11. A method as claimed in claim 10 in which the antigen is selected from
- 12. A method as claimed in claim 11 in which the antigen is the measles H protein. ຜ
- A method as claimed in any one of claims 1 to 12 in which the DNA encoding the antigen is administered only once to the subject. 13,
- encoding the antigen is administered to the subject on at least two occasions. A method as claimed in any one of claims 1 to 12 in which the DNA

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- composition comprising transgenic material is orally administered only once 15. A method as claimed in any one of claims 1 to 14 in which the to the subject. 12
- composition comprising transgenic material is orally administered to the 16. A method as claimed in any one of claims 1 to 14 in which the subject on at least two occasions. ន
- 17. A transgenic plant, the plant having been transformed with a DNA molecule, the DNA molecule comprising a sequence encoding a measles virus andgen such that the plant expresses the measles virus andgen.
- A transgenic plant as claimed in claim 17 in the DNA molecule encodes a fusion protein. 18.

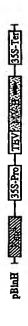
- comprises the measles antigen C-terminally fused to the amino acid sequence 19. A transgenic plant as claimed in claim 18 in which the fusion protein SEKOEL, ဓ္က
- 20. A transgenic plant as claimed in any one of claims 17 to 19 in which the measles antigen is the measles H protein.

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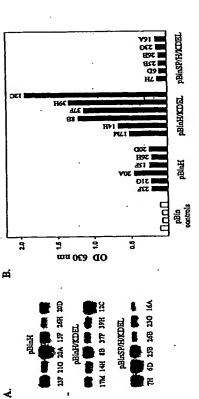
Figure 1



PBINBIKOEL HISBERT - 355-Pro TBV/2002/97-8-3600 - 355-Te-H

PBInSP/H/KDEL H

Figure 2



BINSPAHACDEL

Biatoro

T, transgenic lines

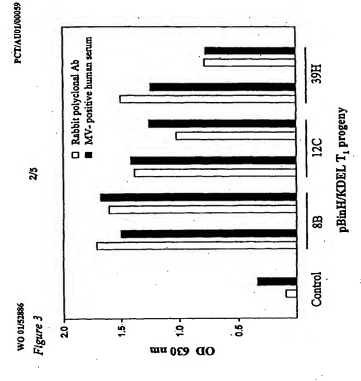
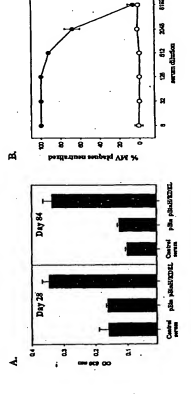
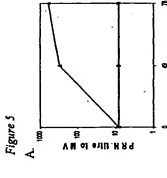


Figure 4



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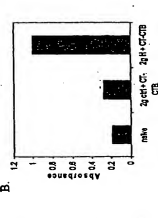


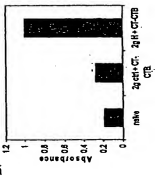


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Figure 7



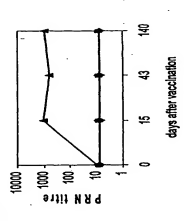


DNA vaccine Plant	Plant feeding		Serum IgG titre	iltre
-			day 0 pre-boost (day 21) post-boost (day 4	post-boost (day 4
MV-H	H	,	•	7290
ပြ	Control	Ŝ	810	2430
control	н		,	270
ပြ	Control	3	\$	8

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days after initial feeding





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Figure 8

		21 (produced) days offer OVA Disk receivable
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		Î
		() () () () () () () () () ()
		21 (pro-bosed) 45 (post- days after man-targe lad DMA vasadination

DNA vaccine	Plant feeding PRN titre	PRN dtr		
		day 0	pre-boost	post-boost
H-vm	н		•	2550
	Control		1150	450
control	Н	Ĭ	·	100
	Control	×	20	80

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SEQUENCE LISTING

7/2

<110> Alfred Hospital Commonwealth Scientific and Industrial Research Organisation The University of Melbourne Australian National University

<120> Prime-boost vaccination strategy

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<170> Patentin Ver. 2.1 12

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PCT/AU01/00059

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protoin fused with ER retention signal 13

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Description of Artificial Sequence: PCR primer \$

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34

INTERNATIONAL SEARCH REPORT

International application No. PCT/AU01/00059

CLASSIFICATION OF SUBJECT MATTER

A61K 39/165; 35/78, A01H 5/00, A61K 48/00

... D H

According to International Patent Classification (IPC) or to both national classification and IPC

FIELDS SEARCHED

dinimum documentation searched (classification system followed by classification symbols)

FILE WPAT AND KEYWORDS BELOW

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FILE MEDLINE

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

DERWENT WPAT and MEDLINE KEYWORDS; measles, measles ()H() protein, antigen, transgenic() plant, immuns, vaccine and DNA or gene Biectronic data base consulted during the international search (name of data base and, where practicable, search terms used)

Relevant to claim No. 1-20 នុ Cardoso A I et al. "Immunization with Plasmid DNA Encoding for the Measles Virus Hemagglutinin and Nucleoprotein leads to Humoral and Cell-Mediated Immunity." Virology, Vol. 225, (1996) Pg 293-299. Citation of document, with indication, where appropriate, of the relevant passages Mahom B P et al. "Approaches to New Vaccines." Clinical Reviews in Biotechnology, Vol. 18(4), (1998), Pg 257-282.
See whole document DOCUMENTS CONSIDERED TO BE RELEVANT See whole document Category × ×

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combination being obvious to a person skilled in the art

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inventive step when the document is taken alone

priority data and not in conflict with the application but cited to understand the principle or theory underlying the invention

Date of spailing of the international scatch report 3 APR 2001

Date of the actual completion of the international search AUSTRALLAN PATENT OFFICE
PO BOX 200, WODEN ACT 2606, AUSTRALIA
B-rnall achress: per@fpsurtvalls.gov.su
Fscstmile No. (02) 6283 3929 Name and mailing address of the ISA/AU 16 March 2001

Meti Sardans ARATI SARDANA

Telephone No: (02) 6283 2627

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ion No.		Rolevant to claim No.	1-20	1-20							 	
International application No.). DOCUMENTS CONSIDERED TO BE RELEVANT	of the relevant passages	body Response to Foot and Mouth Immunization with Alfalfa otein VP 1".	April 1999		-						•
INTERNATIONAL SEARCH REPORT		Cintion of document, with indication, where appropriate, of the relovant passages	Wigdonovitz A et al. "Induction of a Protective Antibody Response to Foot and Mouth Disease Virus in Mice Following Oral or Parenteral Immunization with Alfalfa Transgenic Plants Expressing the Viral Structural Protein VP 1". Virology, Vol. 255, (1999), Pg 347-353. See whole document	WO 99/18225 A (LOMA LINDA UNIVERSITY) 15 April 1999 Sec whole document								
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INTERNATIONAL SEARCH REPORT	International applic
Information on patent family members	PCT/AU01/0005

lication No. 059

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Patent Document Cited in South Report	WO 9918225	-
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